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Object-based approach to programming distributed systems*

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SUMMARY

Two kinds of parallel computers exist: those with shared memory and those without. The former are difficult to build but easy to program. The latter are easy to build but difficult to program. In this paper we present a hybrid model that combines the best properties of each by simulating a restricted object-based shared memory on machines that do not share physical memory. In this model, objects can be replicated on multiple machines. An operation that does not change an object can then be done locally, without any network traffic. Update operations can be done using the reliable broadcast protocol described in the paper. We have constructed a prototype system, designed and implemented a new programming language for it, and programmed various applications using it. The model, algorithms, language, applications and performance will be discussed.

1. INTRODUCTION

Massively parallel computing is just around the corner. Applications such as climate prediction, pharmaceutical design and computers that understand human speech demand more computing power than even a supercomputer can deliver. The solution to these computing needs will have to lie in parallel processing. This paper discusses a new approach to writing the software for these parallel systems.

Architecture for parallel systems falls into two rough categories:

- multiprocessors
- multicomputers

The first category, multiprocessors, are defined as those machines that share a common main memory that all CPUs can read and write using direct machine instructions. Figure 1(a) depicts a simple multiprocessor using a single bus.

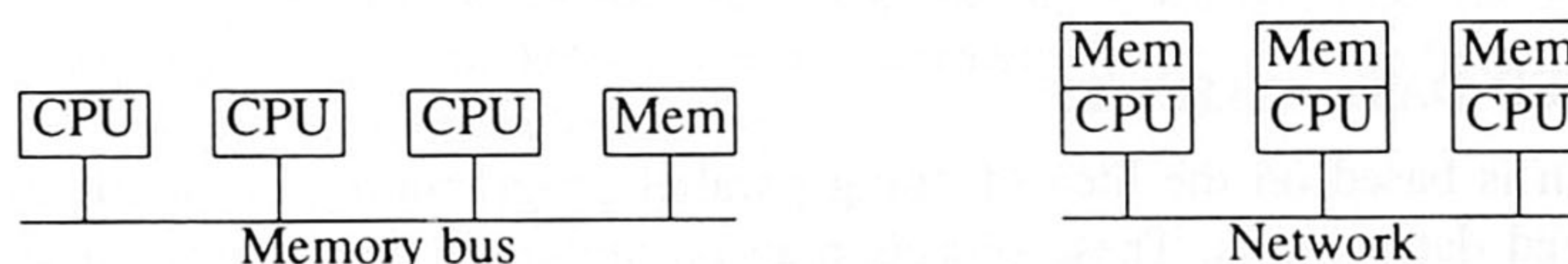


Figure 1. Two kinds of parallel computers: (a) a single-bus multiprocessor; (b) a single-bus multicomputer

*Based on 'Programming a Distributed System Using Shared Objects' by A. S. Tanenbaum, H. E. Bal, S. Ben Hassen and M. Frans Kaashoek which appeared in 2nd International Symposium on High-Performance Distributed Computing (HPDC-2), Spokane, WA, USA, July 1993: pp. 5–12. ©1993 IEEE.

In contrast, multicomputers do not share any main memory. Each one has its own private memory, as shown in Figure 1(b). The CPUs in a multicomputer communicate by sending messages over a network.

Multiprocessors are programmed using well known techniques. Processes on different machines can share variables, synchronizing using semaphores, monitors, critical regions and other simple constructs.

Multicomputers, in contrast, are much harder to program. Typically programs use SEND and RECEIVE primitives to exchange messages. In this way, I/O becomes a central concept in parallel programming. Experience shows that programmers often have difficulties writing error-free parallel programs based on I/O because the level of abstraction is low.

Researchers have been aware of this problem for years. In their pioneering work on IVY, Li and Hudak[1] proposed a solution in which a single virtual address space is shared among a collection of workstations on a network, effectively creating a virtual multiprocessor. Pages can be located on any machine. When a non-local page is referenced, a page fault occurs, and the page handler fetches the page from the machine currently holding it, instead of from disk, as in a traditional virtual memory system. Various optimizations are possible, such as replicating read-only pages, but the performance is frequently too low for practical use.

The next step in this direction is to realize that much of the inefficiency comes from transporting data over the network in fixed-size pages. This can be attacked by making software-defined data structures the unit of sharing, rather than pages. In Linda[2], for example, the virtual shared memory consists of abstract tuples. Processes on any machine can put tuples into the tuple space or take tuples out of the tuple space. Adding or deleting a tuple only requires transporting at most one tuple, not a whole page.

Another approach is taken by Emerald[3], in which the shared memory consists not of pages or tuples, but of abstract data types. Processes can perform operations on abstract data types without regard to where they are located. These operations can be implemented by either moving the request to the abstract data type or by moving the abstract data type to the request.

Both tuple-based and abstract data type-based schemes eliminate the problem found in Ivy and similar systems of having to move fixed-size units (e.g. 8K pages) around, but they have other problems. Emerald does not replicate data, which can lead to performance problems; Linda has fixed primitives that are low-level and inflexible.

2. SHARED DATA-OBJECTS

Our design is based on the idea of doing parallel programming on distributed systems using shared data-objects. These objects may be replicated on multiple processors, and are kept synchronized by system software, the runtime system, as shown in Figure 2. Associated with each shared object is a set of operations that are encapsulated with the object to form an abstract data type.

Processes on different machines can perform operations on shared objects as though they were in physical shared memory. Shared objects exhibit the property of sequential consistency in that if processes simultaneously perform operations on a shared object, the final result will look like all the operations were performed in some sequential order[4]. The order is chosen non-deterministically, but all processes will see the same order of events and the same final result. It is up to the runtime system to maintain this illusion.

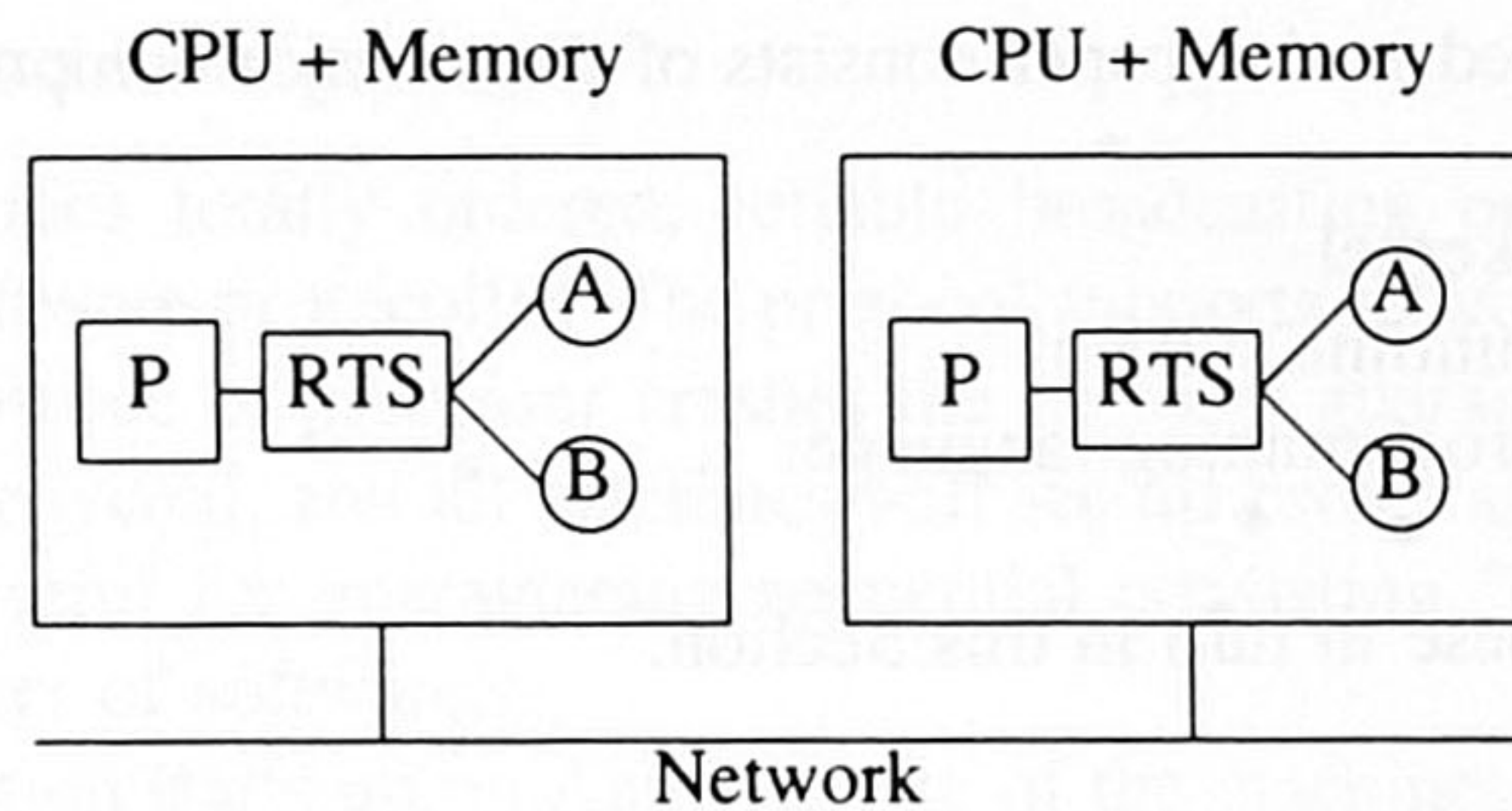


Figure 2. An object can be replicated on each machine: *P* is a process, *RTS* is the runtime system, *A* and *B* are shared data-objects; to the user processes, *A* and *B* look like they are in physical shared memory

By encapsulating the data inside abstract data types, we ensure that processes may not access shared data without the runtime system gaining control. Getting control is essential to make sure objects are consistent when accessed and to guarantee that updates are propagated to other machines in a consistent manner. These properties are not available with a page-based distributed shared memory in which any process can touch any virtual address at will.

Replicating shared objects has two advantages over systems like Ivy. Firstly, reads to any object can be done locally on any machine having a replica. No network traffic is generated. (For our purposes, a read is an operation that does not change the state of its object.) Secondly, more parallelism is possible on reads, since multiple machines can be reading an object at the same time, even if the object is writable. With page-based schemes, having many copies of writable pages is possible only under restricted circumstances involving weakened consistency.

Whether replication can be done efficiently in software depends on two factors. The first is the ratio of reads to writes. If the vast majority of accesses to shared data are reads, then having a copy of each shared object on each machine that needs it is a good idea. The gain from making reads cheap generally results in a major gain in performance. The other factor is how expensive writes are. If writes are exceedingly expensive (in terms of delay, bandwidth, or computing power required), even a moderately high ratio of reads to writes may not be enough to make replication worthwhile. We have studied this question in detail and reported on it elsewhere[5, 6].

3. IMPLEMENTATION

The idea of sharing objects on a distributed system stands or falls with the efficiency of its implementation. If it can be implemented efficiently, high performance parallel systems can be built as multicomputers and programmed as multiprocessors, combining the hardware simplicity of the former with the software simplicity of the latter. If it cannot be implemented efficiently, the idea is of little practical value. Our results show that shared objects can be implemented efficiently under certain circumstances, and give good results for a variety of problems.

The system described in this paper consists of three major components:

1. the Amoeba microkernel
2. the shared object runtime system
3. the Orca parallel programming language.

We discuss each of these in turn in this Section.

3.1. The Amoeba microkernel

Amoeba is a distributed operating system consisting of a microkernel and a collection of server processes[7]. Amoeba was designed for a large number of machines, called the processor pool, connected by a (broadcast) network. There are also machines for handling specialized servers, such as the file system, as well as workstations for clients, but the real computing is done on the processor pool machines. A copy of the microkernel runs on all these machines.

The Amoeba microkernel has four primary functions:

1. manage processes and threads
2. provide low-level memory management support
3. handle I/O
4. support transparent, reliable communication.

Let us consider each of these in turn.

Like most operating systems, Amoeba supports the concept of a process. In addition, Amoeba also supports multiple threads of control, or just *threads* for short, within a single address space. A process with one thread is essentially the same as a process in UNIX[®]. Such a process has a single address space, a set of registers, a program counter and a stack.

In parallel applications, a single process can often have multiple threads. These threads can synchronize using semaphores and mutexes to prevent two threads from accessing critical regions or data simultaneously.

The second task of the microkernel is to provide low-level memory management. Threads can allocate and deallocate blocks of memory, called *segments*. These segments can be read and written, and can be mapped into and out of the address space of the process to which the calling thread belongs. To provide maximum communication performance, all segments are memory resident.

The third basic function of the microkernel is to manage I/O devices, handle interrupts, and so on. Device drivers run as threads within the kernel. All other functionality is located in user-space servers and other processes.

The fourth job of the microkernel is to provide the ability for one thread to communicate transparently with another thread, regardless of the nature or location of the two threads. The model used here is remote procedure call (RPC) between a client and a server[8].

All RPCs are from one thread to another. User-to-user, user-to-kernel and kernel-to-kernel communication all occur. When a thread blocks awaiting the reply, other threads in the same process that are not logically blocked may be scheduled and run.

3.1.1. Totally-ordered broadcasting

Amoeba also provides totally ordered, reliable broadcasting on unreliable networks through use of a software protocol[9]. The protocol supports reliable broadcasting, in the sense that in the absence of processor crashes the protocol guarantees that all broadcast messages will be delivered, and all machines will see all broadcasts in exactly the same order, a property useful for guaranteeing sequential consistency. This feature is heavily used by higher layers of software.

When an application starts up on Amoeba, one of the machines (which normally have identical hardware) is selected as *sequencer* (like a committee electing a chairman), as shown in Figure 3. If the sequencer machine subsequently crashes, the remaining members elect a new one.

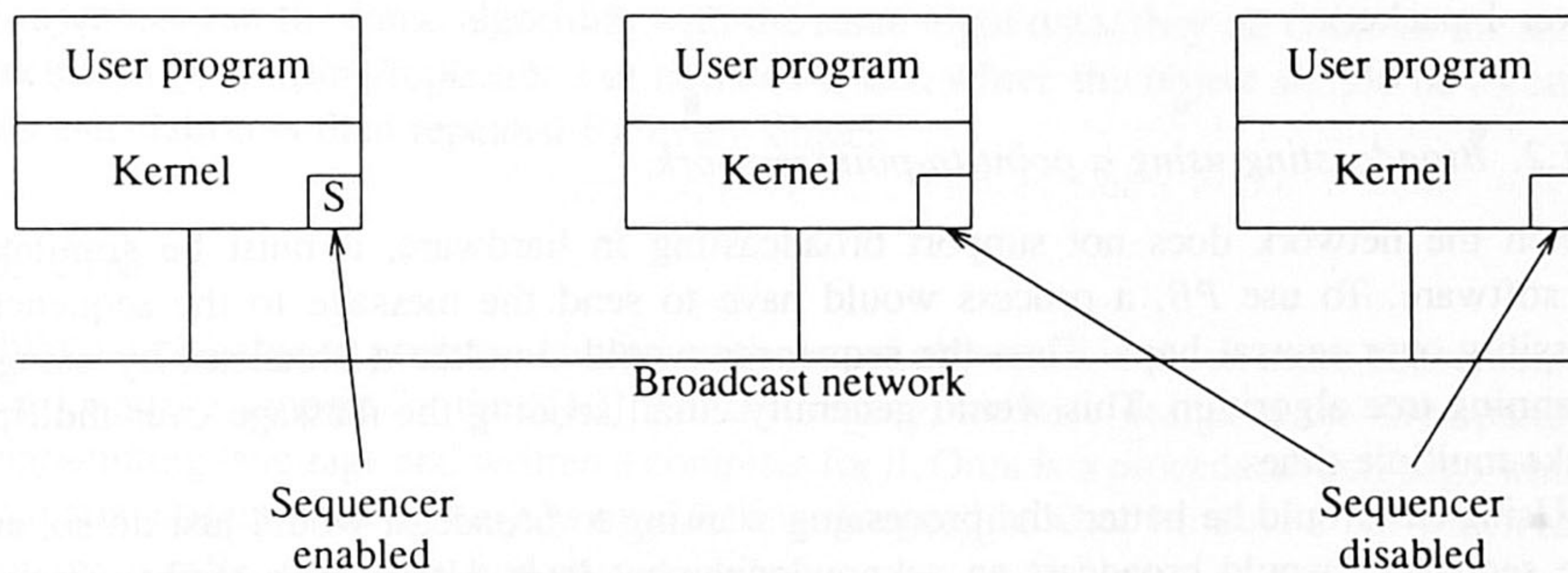


Figure 3. System structure: each kernel is capable of becoming the sequencer, but at any instant only one of them functions as sequencer

We have devised and implemented two different reliable broadcast algorithms having slightly different properties. In the first algorithm, called *PB* (Point-to-point-Broadcast), a runtime system needing to reliably broadcast a message (e.g. a new value of an object) traps to the kernel. The kernel adds a protocol header containing a unique identifier, the number of the last broadcast it has received, and a field saying that this is a *RequestForBroadcast* message and sends it as a point-to-point message to the sequencer. When the sequencer gets the message it adds the lowest unassigned sequence number, stores the message in its history buffer, and then broadcasts it.

When such a broadcast arrives at each of the other machines, a check is made to see if this is the next message in sequence. If this is message 25 and the previous message received was 23, for example, the message is temporarily buffered and a request is sent to the sequencer asking it for message 24 (stored in the sequencer's history buffer). When 24 comes in, 24 and 25 are passed to the application program in that order. Under no circumstances are messages passed to application programs out of order. This is the basic mechanism by which it is guaranteed that all broadcasts are seen by all machines, and in the same order.

The other reliable broadcast algorithm is called *BB* (Broadcast-Broadcast). In this method, the user broadcasts the message, including a unique identifier. When the sequencer sees this, it broadcasts a special *Accept* message containing the unique identifier

and its newly assigned sequence number. A broadcast is only official when the *Accept* message has been sent.

These protocols are logically equivalent, but they have different performance characteristics. In *PB*, each message appears in full on the network twice: once to the sequencer and once from the sequencer. Thus a message of length m bytes consumes $2m$ bytes worth of network bandwidth. However, only the second of these is broadcast, so each user machine is only interrupted once (for the second message).

In *BB*, the full message only appears once on the network, plus a very short *Accept* message from the sequencer, so only half the bandwidth is consumed. On the other hand, every machine is interrupted twice, once for the message and once for the *Accept*. Thus *PB* wastes bandwidth to reduce interrupts compared to *BB*. The present implementation looks at each message and depending on the amount of data to be sent, dynamically chooses either *PB* or *BB*, using the former for short messages and the latter for long ones (over 1 packet).

3.1.2. Broadcasting using a point-to-point network

When the network does not support broadcasting in hardware, it must be simulated in software. To use *PB*, a process would have to send the message to the sequencer, possibly over several hops. Then the sequencer would simulate a broadcast by using a spanning tree algorithm. This would generally entail sending the message over multiple links multiple times.

Using *BB* would be better: the processing wanting to broadcast would just do so, and the sequencer would broadcast an acknowledgement later. However, a slight variation eliminates the second round of broadcasts. In the actual point-to-point protocol, the process wanting to do a broadcast first does an RPC with the sequencer to get a sequence number, then it broadcasts the message and sequence number using a spanning tree algorithm. This algorithm is more efficient than either *PB* or *BB* on point-to-point networks.

3.2. The shared object runtime system

We have developed several runtime systems for Orca over the years. Below we will describe the one currently being tested. It is based on experience with earlier ones. The new runtime system uses information produced by the compiler to do its job. The compiler distinguishes read operations from write operations on shared objects. Operations that do not modify the object are considered reads; the rest are considered writes.

Based on information from the compiler and its own observations, the runtime system can choose, for each object, to replicate it or to maintain a single copy. If a single copy is maintained, the runtime system must decide where to store it. Full replication is best when many processes share an object that is heavily written and rarely read. Keeping only a single copy is best when few processes use it or it is mostly written rather than read.

For each object, the compiler analyses the source code to make an estimate of the number of reads and writes by each process using it. This information is passed to the runtime system, which then maintains read and write statistics for each shared object as execution proceeds. When a new process is created, the replication algorithm is run to

see if any replicated object should be put into single-copy mode or any single-copy mode object should be replicated. Running the replication algorithm only at process creation time is arbitrary, but convenient, since process creation is broadcast to all machines. All machines run the same replication algorithm so they all come to the same replication decision.

The replication algorithm minimizes the estimated number of messages. It first computes the estimated number of messages that will be needed if the object is replicated, taking into account whether or not broadcasting is done in hardware (using the totally ordered broadcast protocols described above) or simulated in software. It then computes the machine making the most total references to the object and computes the number of messages that will be needed if only one copy is maintained, but on the machine making the most references. All other machines must then access the object using RPC. Whichever computation gives the lowest number of messages is chosen. Since all machines run the same algorithm with the same input data, they all come to the same conclusion concerning replication or not, and if not, where the object should be located. This calculation is then repeated for every object.

3.3. Orca

While it is possible to program directly with shared objects, it is much more convenient to have language support for them[10]. For this reason, we have designed the Orca parallel programming language and written a compiler for it. Orca is a procedural language whose sequential statements are roughly similar to languages like C or Modula 2 but which also supports parallel processes and shared objects.

There are four guiding principles behind the Orca design:

- transparency
- semantic simplicity
- sequential consistency
- efficiency

By *transparency* we mean that programs (and programmers) should not be aware of where objects reside. Location management should be fully automatic. Furthermore, the programmer should not be aware of whether the program is running on a machine with physical shared memory or one with disjoint memories. The same program should run on both, unlike nearly all other languages for parallel programming, which are aimed at either one or the other, but not both. (Of course one can always simulate message passing on a multiprocessor, but this is often less than optimal, especially if there is heavy contention for locks and certain other locations.)

Semantic simplicity means that programmers should be able to form a simple mental model of how the shared memory works. Incoherent memory, in which reads to shared data sometimes return good values and sometimes stale (incorrect) ones, is ruled out by this principle.

Sequential consistency is an issue because in a parallel system many events happen simultaneously. By making operations sequentially consistent, we guarantee that operations on objects are indivisible (i.e. atomic), and that the observed behavior is the same as some sequential execution would have been. Operations on objects are guaranteed

not to be interleaved, which contributes to semantic simplicity, as does the fact that all machines are guaranteed to see exactly the same sequence of serial events. Thus the programmer's model is that the system supports operations. These may be invoked at any moment, but if any invocation would conflict with an operation currently taking place, one operation will not begin until the other one has completed.

Finally, *efficiency* is also important, since we are proposing a system that can actually be used for solving real problems.

Now let us look at the principal aspects of Orca that relate to parallelism and shared objects. Parallelism is based on two orthogonal concepts: *processes* and *objects*. Processes are active entities that execute programs. They can be created and destroyed dynamically. It is possible to read in an integer, n , then execute a loop n times, creating a new process on each iteration. Thus the number of processes is not fixed at compile time, but is determined during execution.

The Orca construct for creating a new process is the

fork procname(param, ...)

statement, which creates a new process running the process *procname* with the specified parameters. The user may specify which processor to use, or use the standard default case of running it on the current processor. Objects may be passed as parameters (call by reference). A process may fork many times, passing the same objects to each of the children. This is how objects come to be shared among a collection of processes. There are no global objects in Orca.

Objects are passive. They do not contain processes or other active elements. Each object contains some data structures, along with definitions of one or more operations that use the data structures. The operations are defined by Orca procedures written by the programmer. These operations may contain guards that prevent an operation from starting until certain conditions are met. An object has a specification part and an implementation part, similar in this respect to Ada[®] packages or Modula-2 modules. Orca is what is technically called *object based* (in contrast with *object oriented*) in that it supports encapsulated abstract data types, but without inheritance.

A common way of programming in Orca is the Replicated Worker Paradigm[2]. In this model, the main program starts out by creating a large number of identical worker processes, each getting the same objects as parameters, so they are shared among all the workers. Once the initialization phase is completed, the system consists of the main process, along with some number of identical worker processes, all of which share some objects. Processes can perform operations on any of their objects whenever they want to, without having to worry about all the mechanics of how many copies are stored and where, how updates take place, and so on. As far as the programmer is concerned, all the objects are effectively located in one big shared memory somewhere, but are protected by a kind of monitor that prevents multiple updates to an object at the same time.

4. APPLICATIONS

In this Section we discuss our experiences in implementing various applications in Orca. For each application we describe the parallel algorithm and the shared objects used by the Orca program. In this way, we hope to give some insight into the usefulness of shared objects.

In addition, we briefly consider performance issues of the applications and give some experimental performance results. The performance measurements were carried out on the Amoeba-based Orca system described in the preceding Section using a slightly earlier version of the broadcast runtime system and full replication only. The hardware we use is a collection of MC68030s connected by a 10 Mbit/s Ethernet.

The applications we will look at are: training neural networks, the arc consistency problem, computer chess and automatic test pattern generation. Additional Orca applications are described in [5].

4.1. Training neural networks

A neural network crudely approximates the operation of the human brain. It has neurons and weighted connections between any two neurons (modeling synapses). There are three types of neurons: *input*, *output* and *hidden* neurons. The connections can be *inhibitory* (negative weight) or *excitatory* (positive weight). All connections are unidirectional.

A neuron has two possible states: *off* and *on*, sometimes denoted by 0 and 1, respectively. The activation of a neuron is the weighted sum of the state of the neurons it is connected to. If the activation is above a certain threshold, the neuron is turned on. Otherwise, it is turned off. Each input neuron is externally forced into one of its two states. The others then repeatedly compute their activation until all neuron states are stable. The result is read on the output neurons.

Neurons can be connected in many ways. For instance, a feedforward network consists of at least two layers: an input layer (layer 0), to which the input patterns are applied, and an output layer (layer L). Between the input and the output layers, there may be an arbitrary number of hidden layers. There are connections from each neuron in layer l to all neurons in layer $l + 1$. Neurons within the same layer are not connected to each other, and there are no connections from neurons in a layer to neurons in the preceding layer.

A feedforward network can be trained to recognize an arbitrary set of input-output patterns by modifying the weights on the connections using the backpropagation learning algorithm [11]. Briefly, the learning algorithm works as follows. Start with a random set of weights. For each input pattern, apply the pattern to the network and compute the difference between the desired output and the actual output. Propagate the difference between these two back to the preceding layer and compute the amount with which the connections between layers L and $L - 1$ should be updated using the algorithm given in [11]. Repeat for layer $L - 1$, $L - 2$, and so on, until layer 0 is reached. Now update the weights. This process is repeated for every pattern. The entire set of patterns is applied as many times as necessary, until the network is trained to recognize them.

We have written a parallel implementation in Orca for a slight variation of the algorithm. Instead of updating the weights after each pattern is applied to the network, the update is delayed until all patterns are applied once to the network. The parallelization of the learning process is then very simple: the patterns can be taught in parallel to the network.

Our implementation uses a *network* object, which stores the weights, and two synchronization objects. The patterns are divided into distinct subsets and a subset is

assigned to one teacher process. Thus, there are at most as many teacher processes as input-output pairs. An outline of the teacher process is shown below:

```

while "the network is not trained" do
  phase 1:
    compute the differences for a subset of the patterns;
    wait for the other teachers to finish phase 1;

  phase 2:
    update the weights;
    wait for the other teachers to finish phase 2;
od;

```

Operation *UpdateWeights(WeightDifferences)* of the Network object updates the weights atomically. The weight changes computed by one process are reliably broadcast to all other teacher processes. Because the addition is a commutative operation, the updates for one iteration, from different teacher processes, can be done in any order. Furthermore, this works fine for any number of teacher processes. However, the updates from two different iterations must be serialized. The synchronization objects implement barriers to synchronize the teachers at the end of each phase. At the end of phase 2, the teacher processes must also agree on whether all the patterns are then known by the network or not.

With this implementation, we have obtained reasonable speed-ups. As an example we have experimented with a network of three layers, with ten neurons in each layer. The network was taught 120 input-output pairs. The number of processors was chosen to be a whole fraction of the number of patterns. The results are shown in Figure 4.

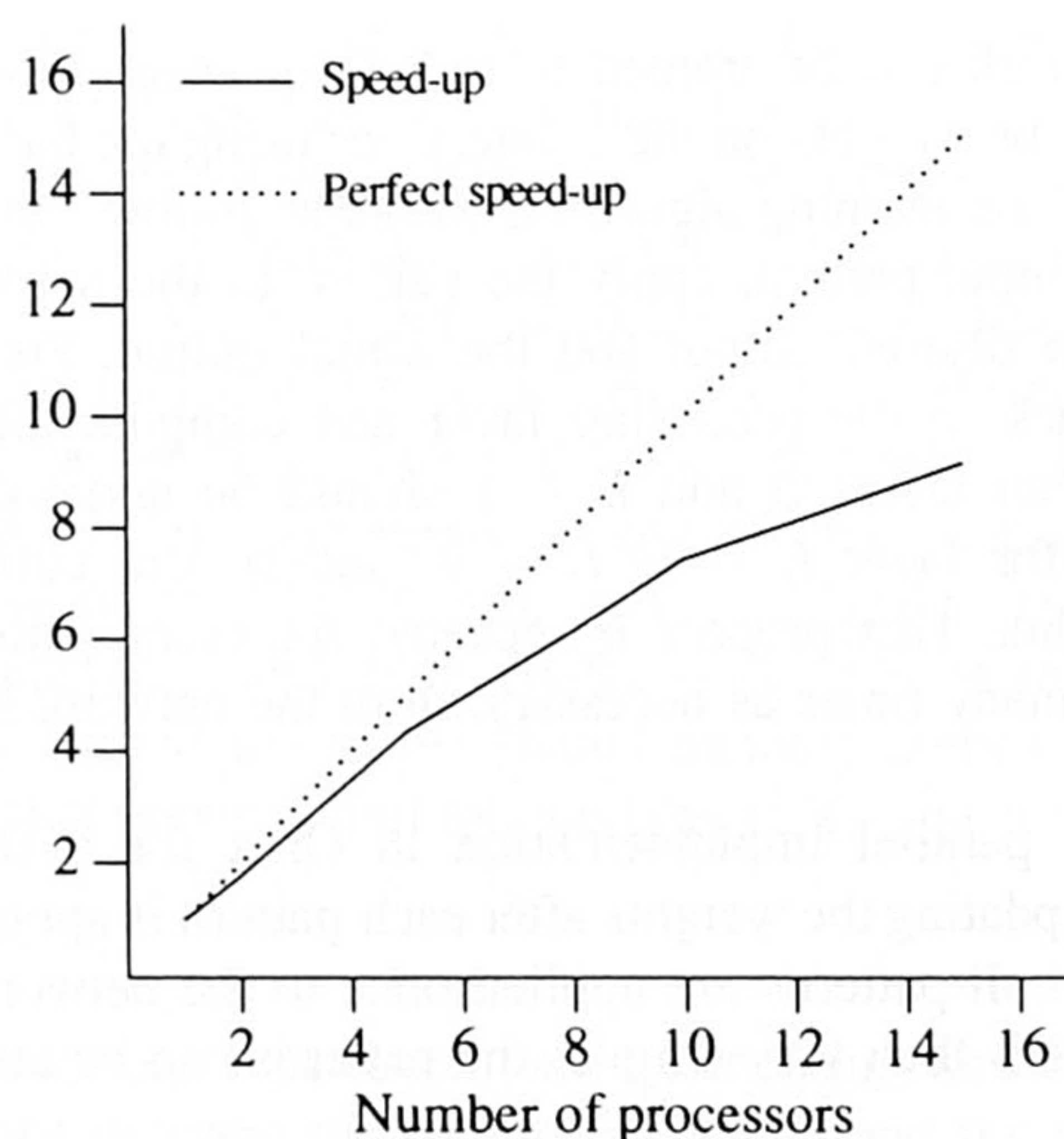


Figure 4. Speed-up for the parallel training of a neural network

4.2. The arc consistency problem

The arc consistency problem (ACP) is an important AI problem[12]. The input to the problem is a set of *variables* V_i , each of which can take a value from a domain D_i , and a list of *constraints*. Each constraint involves two variables and puts a restriction on the values these variables can have, for example $A < B$. The goal of ACP is to determine the maximal set of values each variable can take, such that all constraints are satisfied.

A straightforward sequential algorithm for solving ACP is as follows. Assign a set of possible values to each variable V_i ; initially, the set for V_i contains all values in its domain D_i . Next, repeatedly restrict the sets using the constraints, until no more changes can be made. An obvious improvement is to keep a list of variables whose sets have been changed, and then only recheck constraints involving such variables.

For example, assume the current set for A is $\{1, 10, 100\}$ and the set for B is $\{2, 3, 20\}$. The constraint $A < B$ can be used to delete the value 100 from A 's set. Now, all other constraints involving A have to be checked again.

A parallel implementation of the above algorithm is described by Conrad[13], using a message passing program that runs on an iPSC/2 hypercube. The parallel algorithm statically partitions the variables among the available processors. Each processor takes care of determining the value sets for the variables assigned to it.

We have implemented a similar program in Orca, but now using shared objects, and running on the Ethernet-based Amoeba system. The Orca program uses several shared objects. The sets associated with the variables are stored in a shared object, called a *domain*. This object thus contains an array of sets, one for each variable. Operations exist for initializing the object, deleting an element from one of the sets, and performing set membership tests. The object is shared among all processes, since they all need to have this information.

Another object, called *work*, is used to keep track of which variables have to be rechecked. This object contains an array of Booleans, one per variable. If the value set of a variable A has been checked the entry for A is set to *false*. In addition, for all other variables X for which a constraint exists involving both A and X , the corresponding entry is also set to *true* if A was reduced.

The most complicated issue in parallel ACP is how to terminate the algorithm correctly. The algorithm should terminate if none of the variables needs to be rechecked. Since the variables are distributed among the processors, however, testing this condition requires careful synchronization.

For this purpose, we use two shared objects. One object contains a Boolean variable that is set to *true* if a process discovers that no solution to the input problem exists, because one of its variables now has an empty set of values. Each process reads the object before doing new work, and quits if the value is *true*.

A second and more complicated object, called *result*, contains an array of Booleans, one per process. A process sets its entry to *true* if it is willing to terminate because it has no more work to do. The program can terminate if two conditions are satisfied: (1) all entries in the *work* object are *false*, and (2) all entries in the *result* object are *true*. In this case, no more work exists and neither will any process generate such work, so the program can safely terminate. The *work* and *result* objects have indivisible operations for testing these two conditions.

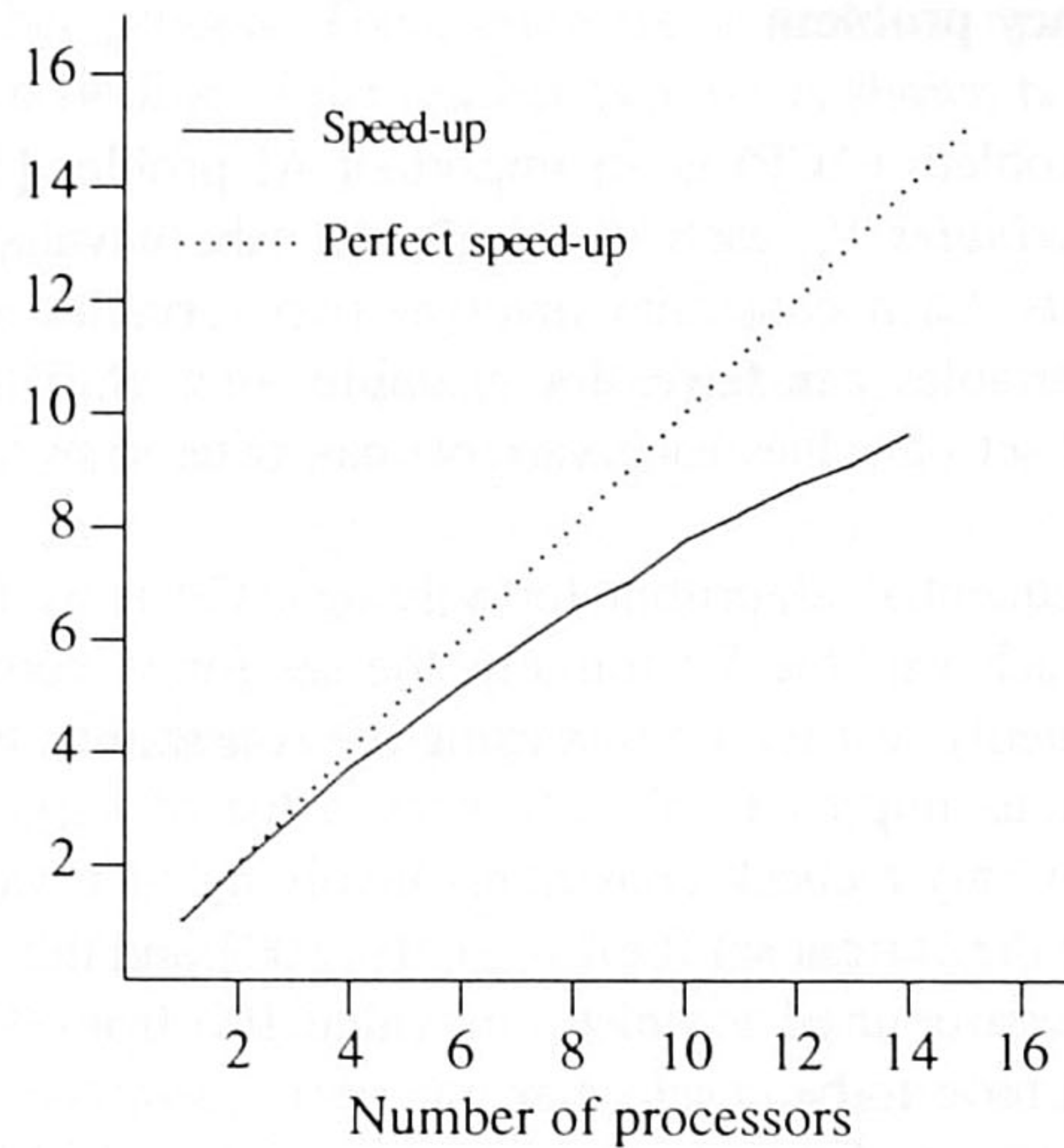


Figure 5. Speed-up for the arc consistency problem using an input problem with 64 variables; one processor was dedicated to work distribution

The speed-ups for ACP are shown in Figure 5. The program uses at least two processors, since the master process that distributes the work runs on a separate processor.

Although the program obtains significant speed-ups, the speed-ups are less than those reported for the original hypercube program. The objects used in the program are replicated on all processors, so there is a lot of CPU overhead in handling incoming update messages for these objects. We should also point out, however, that our program uses many operations to handle termination correctly, as explained above. The hypercube program uses a less expensive but also less elegant method to handle termination, based on time-outs.

4.3. Computer chess

Oracol is a chess problem solver written in Orca. It can be asked to look for 'mate-in- N -moves' or for tactical combinations that win material. It does not consider positional characteristics.

Oracol's search algorithm is based on alpha-beta with iterative deepening and the quiescence search heuristic. Parallelism is obtained by dynamically partitioning the search tree among the different processors, using simple run-time heuristics. The program uses several shared objects (e.g. a job queue). We only discuss two objects here, which are of particular interest.

The two objects implement a *killer table* and a *transposition table*. The killer table contains moves that turn out to cause many cutoffs in the alpha-beta algorithm. Killer moves are always considered first. The idea is that if, say, White threatens to capture a rook by playing 'Queen to a8', then Black needs to do something against this threat. Any move by Black that does not prevent White from capturing the rook can immediately be refuted by the 'Qa8' move, thus saving the trouble of much further analysis.

A transposition table is a table containing positions that have already been analysed earlier during the search. The same board position can be encountered multiple times during the search, because different sequences of moves can result in the same position. The transposition table thus remembers positions and their evaluation values. Before a position is analysed, the program first looks in its transposition table (using a hashing function), to determine if it has seen the position before.

Both the killer table and the transposition table can be implemented as local data structures or as shared objects. If used locally, no communication is needed, but processes cannot benefit from each other's tables. For example, a process may evaluate a position that has been evaluated before by another process. If the tables are shared, this will generally not happen, but now communication overhead is introduced for managing the shared tables.

In Orca, it is particularly easy to implement both versions and see which one is best. The tables are implemented using abstract data types (objects types). In the local version, each process declares its own instance of this type. In the shared version, only the main process declares a table object and passes it as a shared parameter to all other processes. The two versions differ in only a few lines of code. For Oracol, we have determined that, especially for the killer table, shared tables are most efficient.

The speed-ups obtained for the program are not very high, because alpha-beta is hard to parallelize efficiently. On ten CPUs we have measured speed-ups between 4.5 and 5.5. Almost all of the overhead is search overhead, which means that the parallel program searches far more nodes than a sequential program does.

4.4. Automatic test pattern generation

The largest program implemented in Orca so far (nearly 4000 lines) is for automatic test pattern generation (ATPG). ATPG is an important problem from electrical engineering. It generates test patterns for combinatorial circuits. Such a circuit consists of several input and output lines, and several internal gates (e.g. AND gates, OR gates). The output of a given circuit is completely determined by the input.

To test if a specific gate works correctly, the input lines must be set to certain values, such that the correct functioning of the gate can be determined from the output of the circuit. In other words, at least one of the output lines must be different for a correct and an incorrect gate. The problem is how to determine which inputs to use. In general, all gates of the circuit must be tested, so the problem becomes that of generating a set of input patterns that together test the whole circuit. This problem is called the ATPG problem. The problem is NP complete, so in practice an ATPG program tries to cover as many gates as possible within the time limit imposed on it.

Many ATPG algorithms exist, and several parallel algorithms have been designed and implemented[14]. The Orca ATPG program is based on the PODEM algorithm[15]. The algorithm considers each gate in turn. It assigns values to certain input lines (determined by heuristic rules), and propagates these values through the circuit. If it discovers that the current assignment cannot lead to a test of the gate, it backtracks and tries alternative assignments.

The Orca program parallelizes ATPG by statically partitioning the fault set among the processors. Each processor is given a fixed number of gates, for which it computes the

test patterns. Using this basic algorithm, the program achieves good speed-ups (close to linear) on circuits of reasonably large size.

An important optimization that can be applied to both the sequential and parallel ATPG algorithms is *fault simulation*. If a test pattern has been computed for a certain gate, this pattern will probably test other gates in the circuit as well. Fault simulation determines such gates and removes them from the list of gates for which patterns still have to be computed.

This optimization was easy to add to the Orca program. All processes share an object containing the gates for which test patterns have been generated. If a process adds an element to this set, all other processes also use it to determine which gates they can delete from their set. The Orca program using this optimization is faster in absolute speed (by about a factor of 3), but it obtains inferior speed-ups. This is partly due to the communication overhead, and partly to the fact that the static partitioning of work may now lead to a load balancing problem. We intend to use a more dynamic work distribution strategy in the future.

5. SUMMARY

Shared objects offer the possibility of programming certain parallel applications on systems lacking physical shared memory. They offer a model comparable to what programmers of multiprocessors get to see. We believe that the shared object model allows systems to be built that have the ease of construction of multicomputers, combined with the ease of use of multiprocessors. For this reason, we see this model as a promising area for future research.

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